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         AUG 30
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         SEP 25
                 CAS REGISTRY(SM) no longer includes Concord 3D coordinates
         SEP 25
                 CAS REGISTRY(SM) updated with amino acid codes for pyrrolysine
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NEWS 10
         SEP 28
                 CEABA-VTB classification code fields reloaded with new
                 classification scheme
         OCT 19
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                 E-mail format enhanced
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         OCT 23
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                 CAS Registry Number crossover limit increased to 300,000 in
                 multiple databases
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                 The Derwent World Patents Index suite of databases on STN
                 has been enhanced and reloaded
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         OCT 30
                 CHEMLIST enhanced with new search and display field
NEWS 17
         NOV 03
                 JAPIO enhanced with IPC 8 features and functionality
NEWS 18
         NOV 10
                 CA/CAplus F-Term thesaurus enhanced
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        NOV 10
                 STN Express with Discover! free maintenance release Version
                 8.01c now available
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         NOV 20
                 CAS Registry Number crossover limit increased to 300,000 in
                 additional databases
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         NOV 20
                 CA/CAplus to MARPAT accession number crossover limit increased
                 to 50,000
NEWS 22
        DEC 01
                 CAS REGISTRY updated with new ambiguity codes
NEWS 23
         DEC 11
                 CAS REGISTRY chemical nomenclature enhanced
         DEC 14
NEWS 24
                 WPIDS/WPINDEX/WPIX manual codes updated
NEWS 25
        DEC 14
                 GBFULL and FRFULL enhanced with IPC 8 features and
                 functionality
NEWS 26
         DEC 18
                 CA/CAplus pre-1967 chemical substance index entries enhanced
                 with preparation role
NEWS 27
         DEC 18
                 CA/CAplus patent kind codes updated
NEWS 28
         DEC 18
                 MARPAT to CA/CAplus accession number crossover limit increased
                 to 50,000
                 MEDLINE updated in preparation for 2007 reload
NEWS 29
        DEC 18
NEWS 30
        DEC 27
                 CA/CAplus enhanced with more pre-1907 records
             NOVEMBER 10 CURRENT WINDOWS VERSION IS V8.01c, CURRENT
NEWS EXPRESS
              MACINTOSH VERSION IS V6.0c(ENG) AND V6.0Jc(JP)
              AND CURRENT DISCOVER FILE IS DATED 25 SEPTEMBER 2006.
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=> s cataly? (1) steam (4a) reform? (1) adiabatic?

1354027 CATALY?

210687 STEAM

493 STEAMS

211045 STEAM

(STEAM OR STEAMS)

49974 REFORM?

50628 ADIABATIC?

L1 77 CATALY? (L) STEAM (4A) REFORM? (L) ADIABATIC?

=> s l1 and autothermal?

1372 AUTOTHERMAL?

L2 14 L1 AND AUTOTHERMAL?

=> s l2 and (preheat? or pre (la) heat?)(s) (feedstock or hydrocarbon)

49276 PREHEAT?

210586 PRE

699 PRES

210954 PRE

(PRE OR PRES)

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2376185 HEAT?
         17306 FEEDSTOCK
          8004 FEEDSTOCKS
         22508 FEEDSTOCK
                 (FEEDSTOCK OR FEEDSTOCKS)
        334832 HYDROCARBON
        336188 HYDROCARBONS
        517310 HYDROCARBON
                 (HYDROCARBON OR HYDROCARBONS)
          1549 (PREHEAT? OR PRE (1A) HEAT?)(S) (FEEDSTOCK OR HYDROCARBON)
L3
             0 L2 AND (PREHEAT? OR PRE (1A) HEAT?)(S) (FEEDSTOCK OR HYDROCARBON
=> s 12 and (preheat? or pre (1a) heat?)
         49276 PREHEAT?
        210586 PRE
           699 PRES
        210954 PRE
                 (PRE OR PRES)
       2376185 HEAT?
          4429 PRE (1A) HEAT?
L4
             O L2 AND (PREHEAT? OR PRE (1A) HEAT?)
=> s 12 and monolith
          3953 MONOLITH
          2811 MONOLITHS
          5688 MONOLITH
                 (MONOLITH OR MONOLITHS)
L5
             1 L2 AND MONOLITH
=> d 15 ibib ab
     ANSWER 1 OF 1 CAPLUS COPYRIGHT 2007 ACS on STN
L5
ACCESSION NUMBER:
                         2005:20229 CAPLUS
DOCUMENT NUMBER:
                         142:282673
TITLE:
                         Monolithic structures as alternatives to particulate
                         catalysts for the reforming of hydrocarbons for
                         hydrogen generation
AUTHOR (S):
                         Giroux, Thomas; Hwang, Shinn; Liu, Ye; Ruettinger,
                         Wolfgang; Shore, Lawrence
CORPORATE SOURCE:
                         Engelhard Corporation, Iselin, NJ, 08830, USA
SOURCE: .
                         Applied Catalysis, B: Environmental (2005), 56(1-2),
                         185-200
                         CODEN: ACBEE3; ISSN: 0926-3373
PUBLISHER:
                         Elsevier B.V.
DOCUMENT TYPE:
                         Journal; General Review
LANGUAGE:
                         English
     A review of the current status of developments of monolithic precious
     metal catalysts for use in small-scale hydrogen generation from
     hydrocarbons by reforming, steam reforming,
     water gas shift reaction, and preferential oxidation with minimal carbon
     monoxide formation for use in proton-exchange-membrane fuel cells.
     monolithic catalysts are then used for on-site hydrogen
     generators for industrial sites and for hydrogen fueling stations for fuel
     cell-powered automobiles. The advantages of monoliths or heat
     exchangers were emphasized for each of the unit operations. In reforming,
     data were presented from high-space-velocity catalytic partial
     oxidation, autothermal reforming reactors, and coated metal
     substrate catalysts for steam reforming with
     enhanced heat transfer. Water gas shift reactor vols. can be decreased
     significantly using precious metal monolithic catalysts that can
     endure transient and cyclic start/stop operations. Preferential oxidation
     reactions were also described, for example, for use in nearly
     adiabatic ceramic monoliths or nearly isothermal
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behavior on a heat exchanger.

REFERENCE COUNT: 49 THERE ARE 49 CITED REFERENCES AVAILABLE FOR THIS

RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

=> s 12 not 15

DOCUMENT TYPE:

L6 13 L2 NOT L5

=> d 16 ibib ab 1-13

L6 ANSWER 1 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2006:224547 CAPLUS

DOCUMENT NUMBER: 145:474502

TITLE: Autothermal CFB membrane reformer for

hydrogen production from heptane

AUTHOR(S): Chen, Zhongxiang; Elnashaie, Said S. E. H.

CORPORATE SOURCE: Department of Chemical & Biological Engineering,

University of British Columbia, Vancouver, BC, V6T

1Z4, Can.

SOURCE: World Congress of Chemical Engineering, 7th, Glasgow,

United Kingdom, July 10-14, 2005 (2005),

83958/1-83958/10. Institution of Chemical Engineers:

Rugby, UK.

CODEN: 69HUFZ; ISBN: 0-85295-494-8 Conference; (computer optical disk)

LANGUAGE: English

AB H production by steam reforming of heptane is studied in an autothermal circulating fluidized bed (CFB) membrane reformer. Pseudo-steady-state simulations show that when the catalyst is not regenerated, the Ni reforming catalyst deactivates quickly, especially at high temps. Continuous catalyst

regeneration keeps the catalyst activity high (.apprx.1.0) and autothermal operation for the entire adiabatic

reaction-regeneration process is achievable when the exothermic heat

generated from the catalyst regenerator is sufficient to compensate for the endothermic heat consumed in the riser reformer. For this process autothermal operation is the most efficient. This

type of autothermal operation requires careful optimization of the steam to C (S/C) ratio. This ultra-efficient reforming process also shows bifurcation behavior, dictating tight control. Multiplicity occurs when S/C ratio is 1.444-2.251 mol/mol. The maximum net H yield is quite high approaching 15.6 mol H/mol heptane at the lower steady-state when S/C feed

ratio is close to 1.444 mol/mol. REFERENCE COUNT: 34 THERE

REFERENCE COUNT: 34 THERE ARE 34 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 2 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:1246215 CAPLUS

DOCUMENT NUMBER: 144:314873

TITLE: Catalytic autothermal reforming of diesel

fuel for hydrogen generation in fuel cells AUTHOR(S): Cheekatamarla, Praveen K.; Lane, Alan M.

CORPORATE SOURCE: Department of Chemical Engineering, The University of

Alabama, Tuscaloosa, AL, 35487, USA

SOURCE: Journal of Power Sources (2005), 152, 256-263

CODEN: JPSODZ; ISSN: 0378-7753

PUBLISHER: Elsevier B.V.

DOCUMENT TYPE: Journal LANGUAGE: English

AB Polymer electrolyte membrane (PEM) fuel cells require hydrogen as the fuel source for generating power. Hydrogen can be produced in a fuel processor by the catalytic reforming of hydrocarbons. The objective of this paper is to present an anal. of the autothermal reforming (ATR) of synthetic diesel fuel in an adiabatic reactor using a

Pt/ceria catalyst. ATR combines endothermic steam reforming and exothermic partial oxidation reactions in a single unit. This simple system provides higher efficiency and higher energy d. than other conventional processes. The product composition as a function of the operating variables and the temperature and concentration profile inside

the

reactor were studied. Hydrogen was generated under adiabatic conditions by heating the feed mixture and ATR reactor to only 400 °C in contrast to higher temps. reported in the literature. The stability of the catalyst and its response to the presence of S poison was also investigated.

REFERENCE COUNT:

AUTHOR (S):

PUBLISHER:

THERE ARE 45 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 3 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:689567 CAPLUS

DOCUMENT NUMBER: 143:424550

TITLE: Autothermal CFB membrane reformer for

hydrogen production from heptane Chen, Z.; Elnashaie, S. S. E. H.

CORPORATE SOURCE: Department of Chemical & Biological Engineering,

University of British Columbia, Vancouver, Can.

SOURCE: Chemical Engineering Research and Design (2005),

83(A7), 893-899

CODEN: CERDEE; ISSN: 0263-8762 Institution of Chemical Engineers

DOCUMENT TYPE: Journal LANGUAGE: English

AB Hydrogen production by steam reforming of heptane is

investigated in a novel autothermal circulating fluidized bed

(CFB) membrane reformer. Pseudo-steady-state simulations show that when

the catalyst is not regenerated, the nickel reforming

catalyst deactivates quickly, especially at high temps. Continuous

catalyst regeneration keeps the catalyst activity high (.apprx.1.0) and autothermal operation for the entire

adiabatic reaction-regeneration process is achievable when the

exothermic heat generated from the catalyst regenerator is

sufficient to compensate for the endothermic heat consumed in the riser

reformer. For this process autothermal operation is the most efficient. This type of autothermal operation requires careful

optimization of the steam to carbon ratio. This ultra-efficient reforming

process also shows bifurcation behavior, dictating tight control. Multiplicity occurs when steam to carbon ratio is in the range of 1.444-2.251 mol/mol. The maximum net hydrogen yield is quite high approaching 15.6 mol of hydrogen per mol of heptane fed at the lower

steady-state when steam to carbon feed ratio is close to 1.444 mol/mol.
ERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THE

REFERENCE COUNT: 30 THERE ARE 30 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 4 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2005:654679 CAPLUS

DOCUMENT NUMBER: 143:232535

TITLE: Numerical simulation of ethanol autothermal

reactor for hydrogen production

AUTHOR(S): Buck, Gregory A.; Obara, Hiroyuki

CORPORATE SOURCE: South Dakota Sch. Mines Technol., Rapid City, SD,

5770, USA

SOURCE: Tamagawa Daigaku Kogakubu Kiyo (2005), 40, 41-47

CODEN: TDKOBJ; ISSN: 0371-5981

PUBLISHER: Tamagawa Daigaku Kogakubu

DOCUMENT TYPE: Journal LANGUAGE: English

AB At their present state of development, hydrogen fuel cells have the capability to provide adequate power for a wide range of stationary and

mobile applications. Despite this fact, long-term sustainability of this technol. rests upon the production of hydrogen from renewable sources, such as solar photovoltaic hydrolysis, microbial production in natural biol. processes, or chemical conversion (reforming) of bio-hydrocarbons. Among the techniques under current study, the partial oxidation of alcs. and other hydrocarbon fuels using either catalyzed (CPO) or uncatalyzed (POX) reactions, offers great promise. A suitable combination of total and partial oxidation supports hydrogen production from ethanol in an autothermal reforming process that requires no external addition of energy. Furthermore, the autothermal reforming process conducted in a well insulated reactor, produces temps. that promote addnl. hydrogen production through the endothermic steam reforming and the water-gas shift reactions, which may also be catalyzed or uncatalyzed, with the added benefit of lowered carbon monoxide concus. In this study, an adiabatic, axisym. ethanol reforming reactor was simulated assuming the reactants (air and ethanol C2H5OH) and the products (O2, N2, H2O, CO2, CO and H2) to be in the gaseous state. The gas flows in the reactor were assumed to be turbulent, and the chemical kinetics of a simple four reaction system were assumed to be dominated by the turbulent mixing process. Air flow rates into the reactor were varied for fixed ethanol flow rate to obtain three different levels of oxidation The numerical results for the reacting flow show that hydrogen production is maximum for an oxidation level between that corresponding to total oxidation

and

that corresponding to partial oxidation These findings are in qual. agreement with observations from previous exptl. studies.

L6 ANSWER 5 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:670981 CAPLUS

DOCUMENT NUMBER: 141:352589

TITLE: Steady-state modeling and bifurcation behavior of

circulating fluidized bed membrane

reformer-regenerator for the production of hydrogen

for fuel cells from heptane

AUTHOR(S): Chen, Zhongxiang; Elnashaie, Said S. E. H.

CORPORATE SOURCE: Department of Chemical Engineering, Auburn University,

Auburn, AL, 36849-5127, USA

SOURCE: Chemical Engineering Science (2004), 59(18), 3965-3979

CODEN: CESCAC; ISSN: 0009-2509

PUBLISHER: Elsevier Ltd.

DOCUMENT TYPE: Journal LANGUAGE: English

The production of hydrogen for fuel cells by steam reforming of heptane is investigated in a circulating fluidized bed membrane reformer-regenerator system. Palladium based hydrogen permselective membranes are used for hydrogen removal and dense perovskite oxygen permselective membranes are used for oxygen introduction. A series of pseudo-steady-state simulations show that when the catalyst is not regenerated, the circulating nickel reforming catalyst deactivates quickly and the half catalyst activity life for efficient production of hydrogen is quite short, especially at high temps. Efficient continuous catalyst regeneration can keep the catalyst activity high (.apprx.1.0). With continuous catalyst regeneration, autothermal operation for the entire adiabatic reformer-regenerator system is achievable when the exothermic heat generated from the catalyst regenerator is sufficient to compensate for the endothermic heat consumed in the riser reformer. This type of autothermal operation becomes less likely at high steam to carbon feed ratios. This is due to the fact that carbon deposition rate decreases leading to the decrease of autothermal circulating feed temperature and energy-based hydrogen yield (adiabatic hydrogen yield in autothermal reformer-regenerator system). Multiplicity of the steady states for the reformer is possible for this configuration. With the steam to carbon

feed ratio as the bifurcation parameter, multiplicity occurs between the two bifurcation points 1.444 and 2.251 mol/mol. In this multiplicity region, the energy-based hydrogen yield at the upper steady state with high regenerator output temperature is surprisingly the lowest one, while it is the highest one at the lower steady state with low regenerator output temperature The maximum energy-based hydrogen yield is about 15.58 mol of

per mol of heptane fed at the lower steady-state when steam to carbon feed ratio is very close to the bifurcation value of 1.444 mol/mol. After removing the sweep gas steam by downstream cooling and dehumidification, the product hydrogen from steam reforming of

hydrocarbons can be used for fuel cells with high purity (.apprx.100%). REFERENCE COUNT: THERE ARE 35 CITED REFERENCES AVAILABLE FOR THIS 35 RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 6 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:384129 CAPLUS

DOCUMENT NUMBER: 140:425687

hydrogen

Partial oxidation of light paraffins to synthesis gas TITLE:

in short contact-time reactors

Beretta, Alessandra; Forzatti, Pio AUTHOR (S):

Dipartimento di Chimica, Materiali e Ingegneria CORPORATE SOURCE:

Chimica "G. Natta", Politecnico di Milano, Milan,

20133, Italy

Chemical Engineering Journal (Amsterdam, Netherlands) SOURCE:

(2004), 99(3), 219-226

CODEN: CMEJAJ; ISSN: 1385-8947

PUBLISHER: Elsevier Science B.V.

DOCUMENT TYPE: Journal

LANGUAGE: English

The reaction pathways of ethane and propane partial oxidation to synthesis gas were investigated over a Pt/ $\gamma$ -Al203 and a Rh/ $\alpha$ -Al203 catalyst. An annular reactor was used for this purpose at high space velocities and at temps. below 700 °C in order to avoid homogeneous reactions. Under fuel-rich conditions, the Pt-based catalyst produced CO and H2 at high temps. (>550 °C), while CO2 and H2O were the only reaction products at lower temps. of CO and H2 was consistent with direct oxidation reactions, since contact time had no effect on the product distribution, and secondary reactions ( steam and dry reforming) showed negligible activity. The Rh/Al203 catalyst was also active and selective in the partial oxidation of light hydrocarbons, but in this case the production of hydrogen and CO was strongly dependent on contact time, and steam reforming was important even at short contact times. It was concluded that, over rhodium, both direct and indirect routes were probably involved in the formation of CO and H2. The main difference between the two noble metals thus seemed to be that Pt mainly produced CO and H2 by means of O2 (direct routes), while over Rh the light paraffins were converted to CO and H2 by means of O2 and H2O (direct+indirect routes). This could explain the remarkably different behavior of the two systems when tested in high temperature autothermal reactors (T>700 °C). Under adiabatic conditions, the partial oxidation of light paraffins led to large amts. of gas-phase olefinic products over Pt, whereas high selectivities to synthesis gas were found over Rh. The mechanistic results suggest that this different behavior could be due to the varying capability of Pt and Rh surface reactions to compete with homogeneous reactions.

REFERENCE COUNT: THERE ARE 26 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 7 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:269876 CAPLUS

DOCUMENT NUMBER: 140:289193

TITLE: Process for the production of synthesis gas by the steam reforming of a hydrocarbon feed

INVENTOR(S): Aasberg-Petersen, Kim; Dybkjter, Ib; Christensen,

Peter Seier; Rostrup-Nielsen, Thomas; Erikstrup,

Niels; Hansen, Jetts-Henrik Bak

PATENT ASSIGNEE(S): Den.

SOURCE: U.S. Pat. Appl. Publ., 13 pp.

CODEN: USXXCO

DOCUMENT TYPE:

Patent

LANGUAGE:

English

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATE	APPLICATION NO.	DATE		
VG 0004062707						
US 2004063797	A1	20040401	US 2003-667389	20030923		
EP 1413547	A1	20040428	EP 2003-20673	20030911		
R: AT, BE, CH,	DE, DK	, ES, FR, GB	, GR, IT, LI, LU, NL,	SE, MC, PT,		
IĖ, SI, LT,	LV, FI	, RO, MK, CY	, AL, TR, BG, CZ, EE,	HU, SK		
JP 2004149406	Α	20040527	JP 2003-331066	20030924		
CA 2442491	A1	20040326	CA 2003-2442491	20030925		
NO 2003004285	A	20040329	NO 2003-4285	20030925		
AU 2003248389	A1	20040422	AU 2003-248389	20030925		
ZA 2003007450	Α	20040706	ZA 2003-7450	20030925		
KR 2004027440	Α	20040401	KR 2003-66831	20030926		
CN 1498850	A	20040526	CN 2003-164880	20030926		
PRIORITY APPLN. INFO.:			DK 2002-1435	A 20020926		
AR A process and system	for th	no production	of symthogic and /i			

A process and system for the production of synthesis gas (i.e., H2-CO mixts.) AB from a hydrocarbon feed (e.g., natural gas) comprises endothermic and/or adiabatic catalytic steam reforming and autothermal steam reforming in series, where the steam reforming is carried out in one or more endothermic stages in series or in one or more adiabatic steam reforming stages in series with intermediate heating of the feed stock gas leaving the adiabatic reforming stages and where the carbon monoxide-containing gas, characterized by having a molar ratio of hydrogen to carbon of less than 4.5, is added prior to at least one of the endothermic or adiabatic steam reforming stages and/or prior to the autothermal steam reforming step. Process flow diagrams are presented.

ANSWER 8 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 2004:121:127 CAPLUS

DOCUMENT NUMBER:

140:166150

TITLE:

Hydrogen Production and Carbon Formation during the Steam Reformer of Heptane in a Novel Circulating

Fluidized Bed Membrane Reformer

AUTHOR (S):

Chen, Zhongxiang; Yan, Yibin; Elnashaie, Said S. E. H. Department of Chemical Engineering, Auburn University,

Auburn, AL, 36849-5127, USA

SOURCE:

Industrial & Engineering Chemistry Research (2004),

43(6), 1323-1333

CODEN: IECRED; ISSN: 0888-5885

PUBLISHER:

American Chemical Society

DOCUMENT TYPE:

Journal

LANGUAGE:

CORPORATE SOURCE:

English

Hydrogen production and carbon formation during the steam reforming of heptane over nickel-based catalyst are investigated in a circulating fluidized bed membrane reformer (CFBMR) at 723-823 K and 101.3-2026 kPa. A random carbon deposition and catalyst deactivation model is developed to account for the effect of carbon deposition on the overall reforming kinetics. The results show that the deposited carbon can be efficiently gasified by steam, hydrogen, carbon dioxide, or oxygen in this novel CFBMR, making carbon-free

operation practically possible, especially when steam to carbon feed ratio is higher than 2.5 mol/mol. The use of hydrogen permselective membranes breaks the equilibrium barriers associated with the reversible re-forming reactions and increases the hydrogen yield significantly. introduction of oxygen into the adiabatic reformer can efficiently supply the heat necessary for the endothermic steam reforming through the exothermic oxidation, making an autothermal condition possible for the efficient production of hydrogen.

REFERENCE COUNT:

40 THERE ARE 40 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

ANSWER 9 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

2001:864736 CAPLUS

DOCUMENT NUMBER:

136:8552

TITLE:

Autothermal catalytic steam reforming of

hydrocarbons

INVENTOR(S):

Ahlborn, Rainer; Baumann, Frank; Wieland, Stefan Omg Ag & Co. Kg, Germany; Umicore Ag & Co. KG

PATENT ASSIGNEE(S):

SOURCE:

Eur. Pat. Appl., 9 pp. CODEN: EPXXDW

DOCUMENT TYPE:

Patent

LANGUAGE:

German

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.			KIN	KIND DATE		Α	APPLICATION NO.						DATE					
						_	<b>-</b> -	<del>-</del>	-					<b>-</b>		_		
EP	1157	968			A1		2001	1128	E	P	200	1-1	108	16		2	0010	504
EP	1157	968			В1		2006	0503										
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		ΙE,	SI,	LT,	LV,	FI,	RO,	CY,	TR									
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AT	3250	75			T		2006	0615	A	Т	200	1 - 1	108	16		2	0010	504
US	2002	0094	80		<b>A1</b>		2002	0124	U	S	200	1-8	539	02		2	0010	514
US	7083	775			B2		2006	0801										
CA	2348	120			A1		2001	1120	С	Α	200	1-2	348	120		2	0010	517
JP	2002	0124	80		Α		2002	0115	J	P	200	1-1	476	01		2	0010	517
BR	2001	00204	41		Α		2001	1218	В	R	200	1-2	041			2	0010	518
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AB Autothermal catalytic steam

> reforming is carried out by leading a feed mixture of hydrocarbons, 02, and water or steam over a reforming catalyst. The reaction is operated adiabatically, and the reforming catalyst contains a Pt-group metal deposited on a support from a group of Al2O3, SiO2, TiO3, or their mixed oxide and zeolite. A steam/C ratio in the feed is (0.7-4):1. synthesis gas is suitable for manufacture of H2 for fuel cells.

REFERENCE COUNT:

THERE ARE 6 CITED REFERENCES AVAILABLE FOR THIS ' RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

1.6 ANSWER 10 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

6

ACCESSION NUMBER:

1996:330713 CAPLUS

DOCUMENT NUMBER:

125:14538

TITLE:

Simulation studies of autothermal reactor

system for H2 production from methanol steam reforming

AUTHOR (S):

Ma, L.; Jiang, C.; Adesina, A. A.; Trimm, D. L.;

Wainwright, M. S.

CORPORATE SOURCE:

School of Chemical Engineering and Industrial

Chemistry, University of New South Wales, Sydney, NSW,

Australia

SOURCE:

Chemical Engineering Journal (Lausanne) (1996), 62(2),

103-111

CODEN: CMEJAJ; ISSN: 0300-9467

PUBLISHER: Elsevier DOCUMENT TYPE: Journal LANGUAGE: English

This paper reports the results of a math. investigation on the performance of a class of adiabatic dual-bed catalytic reactor systems with cylindrical and spherical geometries that may be used to promote internal heat exchange for the coupled reaction network. Anal. shows that, while the coaxial cylindrical system and the dual-bed single tubular reactor generally need optimal water-to-methanol feed ratios of about 3-4, the spherical arrangement always requires a ratio <1 for equivalent or even better performance. The spherical reactor system in which the oxidation catalyst was placed in the inner sphere with steam reforming in the annular space showed the most promising performance in terms of reactor efficiency (.apprx.80%) and H2 production (125 m3 gas/m3 reactor/s) while the coaxial reactors exhibited the poorest efficiency (<10%) for a H2 production rate of 19.5 m3 qas/m3 reactor/s. Thus the spherical reactor with an inner oxidation catalyst bed is the most attractive configuration for this autothermal process in terms of product maximization, feed and energy minimization.

ANSWER 11 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER: 1995:452928 CAPLUS

DOCUMENT NUMBER: 122:269347

Reactor modeling and simulations in synthesis gas TITLE:

production

AUTHOR (S): De Groote, Ann M.; Froment, Gilbert F.

CORPORATE SOURCE: Laboratorium voor Petrochemische Techniek,

Universiteit Gent, Ghent, B9000, Belg. SOURCE:

Reviews in Chemical Engineering (1995), 11(2), 145-83

CODEN: RCEGD6; ISSN: 0167-8299

PUBLISHER: Freund DOCUMENT TYPE: Journal LANGUAGE: English

Steam and CO2 reforming, partial oxidation and autothermal reforming processes for synthesis gas production were considered. Detailed kinetics are required for the successful simulation of these processes. Kinetic equations for steam and CO2 reforming were developed based on elementary steps between adsorbed species. Since the reactions are very fast, severe diffusion limitations occur. The thresholds for carbon formation based on methane cracking and the Boudouard reaction were determined in an electrobalance Tubular reactors for steam and CO2 reforming were modeled based on a one-dimensional heterogeneous model accounting for internal diffusion limitations. In these simulations the heat fluxes along the reactor were calculated from a detailed temperature distribution in the

furnace using the FURNACE computer program. Tubular reactors for steam and CO2 reforming were also simulated based on a two-dimensional model accounting for both radial and axial concentration gradients, leading to more accurate bounds for carbon deposition. catalytic partial oxidation of CH4/O mixts. in an adiabatic fixed bed reactor was simulated based on the kinetics for total combustion, steam reforming, and water gas shift reaction. Finally, the catalytic partial oxidation of CH4 to synthesis gas in an adiabatic reversed flow reactor was modeled. This is a transient operation and the traveling waves of temperature and

were simulated for typical operating conditions.

ANSWER 12 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

1993:606619 CAPLUS ACCESSION NUMBER:

DOCUMENT NUMBER: 119:206619

TITLE: Steam reforming. Opportunities and limits of the technology

AUTHOR (S): Rostrup-Nielsen, Jens; Dybkjaer, Ib; Christiansen,

CORPORATE SOURCE:

Haldor Topsoe A/S, Lyngby, DK-2800, Den.

SOURCE:

NATO ASI Series, Series E: Applied Sciences (1992), 225 (Chemical Reactor Technology for Environmentally

Safe Reactors and Products), 249-81

CODEN: NAESDI; ISSN: 0168-132X

DOCUMENT TYPE: Journal; General Review

LANGUAGE: English

A review, with 57 refs., of advances in reactor mech. design and modeling of the conventional tubular reformer, as well as the heat-exchange reformer. Other topics covered include the role of adiabatic pre-reforming, autothermal reforming, CO2 reforming, main requirements of the catalysts, control of tube wall temperature, carbon deposit formation, manufacture of synthesis gas (with examples), and a comparison of direct conversion of CH4 with indirect conversion via steam reforming.

ANSWER 13 OF 13 CAPLUS COPYRIGHT 2007 ACS on STN

ACCESSION NUMBER:

1988:115635 CAPLUS

DOCUMENT NUMBER:

108:115635

TITLE:

Autothermal production of synthesis gas

INVENTOR(S):

Lewis, Jerry Lee

PATENT ASSIGNEE(S):

Fluor Corp., USA

SOURCE:

Faming Zhuanli Shenqing Gongkai Shuomingshu, 9 pp.

CODEN: CNXXEV

DOCUMENT TYPE:

Patent

LANGUAGE:

Chinese

FAMILY ACC. NUM. COUNT:

PATENT INFORMATION:

PATENT NO.	KIND	DATÉ	APPLICATION NO.	DATE
			<b></b>	
CN 85101360	Α	19870131	CN 1985-101360	19850401
CN 1008350	В	19900613		
. השואד זווממג שחדמ			ON 1005 101260	

PRIORITY APPLN. INFO.: CN 1985-101360 19850401 An apparatus for steam reforming of hydrocarbon feedstocks to produce H-rich fuel gases comprises a vertical reactor equipped with >1 reforming tubes packed with catalysts, >1 buffering plates fixed on the reactor walls, and a combusting chamber having an extended pipe communicated with the bottom of the reforming tubes. The hydrocarbon feedstocks are preferably the byproduct effluents (containing MeOH or EtOH) from an NH3-manufacturing plant, mixed with steam, and fed into the top of the reforming reactor at 900-1300°F. The O-rich air is injected from the bottom of the combustion chamber, and the gaseous effluents from the combustion chamber are introduced into a 2nd catalytic reforming zone, and then passed through the outer surface of the reforming tubes. The inner walls of the reforming reactor are preferably made of adiabatic materials to prevent heat loss. The apparatus reduces operation time and increases the yield of H-rich fuel gases.

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**END OF SEARCH HISTORY**